PDBeFold (SSM: Secondary Structure Matching) http://pdbe.org/fold/

This PDBe tutorial introduces the PDBeFold service, an interactive service for comparing protein structures in 3D. This service provides:

- pairwise and multiple comparison and 3D alignment of protein structures
- examination of a protein structure for similarity with the whole Protein Data Bank (PDB) archive or SCOP.
- best Cα-alignment of compared structures
- download and visualisation of best-superposed structures using various graphical packages

PDBefold structure alignment is based on identification of residues occupying "equivalent" geometrical positions. In other words, unlike sequence alignment residue type is neglected. The PDBefold service is a very powerful structure alignment tool which can perform both **pairwise** and **multiple** three dimensional alignment. In addition to this there are various options by which the results of the structural alignment query can be sorted. The results of the Secondary Structure Matching can be sorted based on the Q score (C α -alignment), P score (taking into account RMSD, number of aligned residues, number of gaps, number of matched Secondary Structure Elements and the SSE match score), Z score (based on Gaussian Statistics), RMSD and %Sequence Identity.

It is hoped that at the end of this tutorial users will be able to use PDbeFOLD for the analysis of their own uploaded structures or entries already in the PDB archive.

How to use this tutorial

Please keep this tutorial open in a browser or PDF viewer and then start the tutorial by opening a new instance of your internet browser and follow the steps outlined in the tutorial using that browser instance

Tutorial

PDBeFOLD may be accessed from multiple locations on the PDBe website. From the PDBe home page (http://pdbe.org/), there are two access points for the program as shown below.



The link highlighted in blue may be used to automatically get all related structures to a given PDB code. In order to start the webservice, click on the link shown in red above. This will open up a introductory page concerning PDBeFOLD that contains additional information, tips and help regarding the webservice. On this page, now click on the "Start PDBeFOLD" button.



This should start up the actual webservice that will in the form of a submission form as shown below.

Submission Form for explanation	pairwise 3D alignment multiple on of input
Query	Target
Source: PDB entry : PDB code 1sar view Select chains : (Find chains) Chains: (r(all))	Source: All PDB archive
Lowest acceptable match (%) 70	Lowest acceptable match (%) 70
 ✓ match individual chains ✓ match connectivity ✓ if no matches within acceptibility limit Precision: normal → Sort by: Q-scd 	✓ best matches only ✓ unique matches only ts found, show <i>some</i> of the close ones re ✓ Viewer: ♥
Precision: normal Sort by: Q-sco	Back to Home Page

The form contains parameters that may be adjusted or tweaked in order to customize the results as required. The "Query" contains information regarding the structure/structures that need to be compared. This could be an existing PDB

code, an uploaded coordinate file, SCOP entry or pair of existing PDB entries that need to be compared. In addition to this, if an entry contains more than one chain, the chain that has to be compared can be chosen. Similarly the target for the search could be another PDB entry, uploaded coordinate file or SCOP set or a set of files. The "lowest acceptable match" boxes tell the program what cut-off to use in the matching process. The default is 70% both for the query and target, meaning that in order to list a comparison as a match, at least 70% of the secondary structure of the query must match 70% of the target structure. These settings can be adjusted. Setting this to a lower value will result in a large number of hits and the opposite may result in only identical structures found.

We will use the PDB entry 2MJP (<u>http://pdbe.org/2mjp</u>) (STRUCTURE-BASED IDENTIFICATION OF THE BIOCHEMICAL FUNCTION OF A HYPOTHETICAL PROTEIN FROM METHANOCOCCUS JANNASCHII:MJ0226) as an example for demonstrating PDBefold.



Click on the "Submit your query" button. This will start a alignment job on the farm which will take upto 2-3 minutes depending on server load. Once the results are calculated, a new page showing the same will be displayed on the browser (Similar to that shown below).

			*	Dunad	N.	N	0/														
##	Q	Ρ	z	Rmsa	Naign	Ng	⁷ oseq	%sse	Match	%sse	N _{res}	×	Title								
1	1.00	26.8	15.5	0.00	184	0	100	100	2mjp:A	100	184		STRUCTURE-BASED IDENTIFICATION OF THE BIOCHEMICAL FUNCTION OF A HYPOTHETICAL PROTEIN FROM METHANOCOCCUS JANNASCHIIMJ0226								
2	1.00	25.4	15.0	0.20	184	0	100	100	1b78:A	100	184		STRUCTURE-BASED IDENTIFICATION OF THE BIOCHEMICAL FUNCTION OF A HYPOTHETICAL PROTEIN FROM METHANOCOCCUS JANNASCHILMJ0226								
<u>3</u>	0.95	22.3	14.1	0.49	182	1	99	93	1b78:B	93	184		STRUCTURE-BASED IDENTIFICATION OF THE BIOCHEMICAL FUNCTION OF A HYPOTHETICAL PROTEIN FROM METHANOCOCCUS JANNASCHILMJ0226								
<u>4</u>	0.95	20.0	13.3	0.55	182	1	99	86	2mjp:B	100	184		STRUCTURE-BASED IDENTIFICATION OF THE BIOCHEMICAL FUNCTION OF A HYPOTHETICAL PROTEIN FROM METHANOCOCCUS JANNASCHILMJ0226								
5	0.85	15.3	11.6	1.14	182	2	50	93	2dvn:B	100	186		STRUCTURE OF PH1917 PROTEIN WITH THE COMPLEX OF IMP FROM PYROCOCCUS HORIKOSHII								
6	0.68	9.1	8.9	1.92	181	3	49	93	1v7r:A	100	186		STRUCTURE OF NUCLEOTIDE TRIPHOSPHATE PYROPHOSPHATASE FROM PYROCOCCUS HORIKOSHII OT3								
Ţ	0.67	8.5	8.6	1.96	181	3	49	93	2dvo:A	100	185		STRUCTURE OF PH1917 PROTEIN WITH THE COMPLEX OF ITP FROM PYROCOCCUS HORIKOSHII								
<u>8</u>	0.67	9.0	8.9	1.95	181	3	49	93	2dvn:A	100	186		STRUCTURE OF PH1917 PROTEIN WITH THE COMPLEX OF IMP FROM PYROCOCCUS HORIKOSHII								
9	0.66	6.7	7.7	1.93	177	5	50	79	2dvp:A	92	184		STRUCTURE OF NTPASE FROM PYROCCOUS HORIKOSHII								
<u>10</u>	0.65	7.9	8.3	2.08	181	3	49	93	2e5x:A	100	185		STRUCTURE OF NUCLEOTIDE TRIPHOSPHATE PYROPHOSPHATASE FROM PYROCOCCUS HORIKOSHII 0T3								
11	0.65	8.7	8.8	1.42	168	7	35	71	2car:B	83	194		CRYSTAL STRUCTURE OF HUMAN INOSINE TRIPHOSPHATASE								
12	0.64	8.6	8.8	1.48	170	7	35	71	2car:A	83	196		CRYSTAL STRUCTURE OF HUMAN INOSINE TRIPHOSPHATASE								
13	0.64	8.8	8.8	1.41	168	7	35	71	2i5d:A	83	195		CRYSTAL STRUCTURE OF HUMAN INOSINE TRIPHOSPHATE PYROPHOSPHATASE								
<u>14</u>	0.63	5.4	7.0	1.81	173	6	35	71	1vp2:A	91	189		CRYSTAL STRUCTURE OF PUTATIVE XANTHOSINE TRIPHOSPHATE PYROPHOSPHATASE/HAM1 PROTEIN HOMOLOG (TM0159) FROM THERMOTOGA MARITIMA AT 1.78 A RESOLUTION								

The results here are sorted based on Q-score (Quality of alignment, with 1 being the highest score) for this entry. The pairwise alignment result between **2mjp** and **2e5x** is highlighted in the above figure. There is a **49% amino acid sequence identity** between the two proteins, whereas they have **93% secondary structure identity**. Clicking on the number link on the left hand side of the page will return residue-by-residue description about the structural alignment between the two proteins. *Note: as the PDB archive continuously adds new entries every week, it is possible that this result may not appear in the same position as shown.*

Q	uery P	DB 2mjj	p:A		Alig	gnment		Target PDB 2e5x:A					
N res	%res	N	[%] SSE	Q	Ρ	RMSD	N algn	N _{res}	N _{SSE}	% SSE			
184	98	14	93	0.650	7.93	2.081	181	185	98	13	100		
	STRUCTI	JRE-BAS	ED	0/									
IDE		TION OF	THE ON OF A	~seq	2	SSE	gaps	STRUCTURE OF NUCLEOTIDE					
HYPOT	HETICA	L PROTE	N FROM	48.6	8.35	13	3	PYROPHOSPHATASE FROM					
	METHAN	OCOCCU	JS 26					PYROCOCCUS HORIKOSHII O					
vie	w dow	nload		V	iew su	perposed			view	downlo	bad		
			Viewer:										

Click on the "View Superposed" button to view the secondary structure alignment.



The two structures 2MJP and 2E5X appear share a high level of secondary structure identity despite only having <50% sequence identity. This indicates that the two proteins belong to the same structural family.

A little further down the page are the summary of the alignment showing the secondary structure elements that match between the two structures with residue ranges and the matrix required to move and align the target to the query structure.

	Q	uer	y١	PDB	2mjj	o:A						Т	ar	get F	DB 2	2e5x	: A	
1	SD	4	A	ILE	11	ALA	14	1	<->	1	SD	5	A	LYS	2	ILE	6	
2	H1	11	Α	PRO	18	LEU	28		<->	2	Н1	14	A	ASN	9	THR	22	
3	SD	4	Α	ILE	36	ILE	39		<->	3	SD	5	A	GLU	26	LEU	30	
4	H1	18	Α	THR	49	LYS	66		<->	4	Н1	14	Α	LYS	41	LYS	54	
5	SD	10	Α	VAL	69	VAL	78		<->	5	SD	10	A	PHE	61	ILE	70	
6	H1	9	Α	TYR	88	ILE	96		<->	6	H1	9	A	TYR	80	ILE	88	
7	H1	10	A	ILE	96	LEU	105		<->	7	H1	10	A	ILE	88	MET	97	
8	SD	12	A	ASN	112	ASP	123		<->	8	SD	12	A	ARG	104	ILE	115	
9	SD	13	A	GLY	126	VAL	138		<->	9	SD	13	A	LYS	118	ILE	130	
11	SD	3	A	PHE	155	PRO	157		<->	10	SD	3	A	PHE	146	PRO	148	
12	H5	5	Α	THR	163	MET	167		<->	11	Н5	5	A	THR	154	MET	158	
13	H1	6	A	THR	168	SER	173		<->	12	H1	9	A	THR	159	SER	167	
14	H1	16	A	SER	176	ASP	191		<->	13	H1	18	A	SER	167	LEU	184	
SCO	P: do	oma	in	3322	<u>3</u> , fa	mily g	c.51.4.	1			F	DBe	At	las	PDBe	Motif		<u>\</u>
P	DBe	Atla	S	PDB	e Mot	if I <u>OC</u>	A			Gen	eCer	nsus	I <u>F</u>	SSP	3De	<u>e I CA</u>		PDBsum
GeneCer	nsus	FS	SP	1 <u>3D</u>	<u>ee</u> <u>C</u>	ATH	PDB	sum										
vie	w) (dow	nlo	ad see	quence		(view	superp	osed)		V	iew	dowr	nload s	equen	ce

Rotation-translation matrix

		(to be ap	plied to th	ne t	tar	rget)
Download the page content	-0.586	-0.334	-0.738	1	X		46.884
>> in plain text	-0.714	0.643	0.276	×	Y	+	-30.053
>>) in XML	0.383	0.689	-0.616		Z		-19.129

If you scroll down the page, there is a 3D structural alignment between the residues from the corresponding PDB entries. This alignment shows the extent of superposition between similar secondary structure folds. The red background indicates that matched residues have same residue name, cyan background is used for other matched residues. Unmatched residues are indicated by black background. The letters H and S represent helix and β -strands respectively.

		<u>3D</u> :	Stru	ctural alignm	ner	<u>it</u>				
				notations						
P	DB 2mj	p:A	SI	Dist. (Å)	PDB 2e5x:A					
				·	-	A:MET	1			
+	A:LYS	10	:::•	2.20	S+	A:LYS	2			
s-	A:ILE	11	::•	2.05	s-	A:ILE	3			
s٠	A:TYR	12	::•	1.69	s-	A:PHE	4			
s-	A:PHE	13	::•	1.35	s-	A:PHE	5			
s-	A:ALA	14	:	1.19	s-	A:ILE	6			
•	A:THR	15	::•	1.83	•	A: THR	7			
-	A:GLY	16		2.95	•	A:SER	8			
+	A:ASN	17	:::•	3.82	H+	A:ASN	9			
H+	A:PRO	18	:::	4.87	H+	A:PRO	10			
H+	A:ASN	19		5.09	H–	A:GLY	11			
H+	A:LYS	20	::•	3.70	H+	A:LYS	12			
H-	A:ILE	21	::	3.77	H-	A:VAL	13			
H+	A:LYS	22	::	4.19	H+	A:ARG	14			
H+	A:GLU	23		3.13	H+	A:GLU	15			

The residue-by-residue mapping provides a very useful tool to analyse the structure function relationship between the two entries.

Lets now concentrate on the proteins themselves to get more information about them. You can go to the summary pages for these entries at the PDBe in new browser windows or tabs. Go to http://www.pdbe.org/2mjp in one window and http://www.pdbe.org/2e5x in another. On each of these pages, click on the "Ligands" link from the left-hand sidebar.



As you can see from the above screenshots, both entries contain nucleotide-type ligands. The ligand bound to 2MJP is ANP (PHOSPHOAMINOPHOSPHONIC ACID-ADENYLATE ESTER). For 2E5X, the bound nucleotide is ITT (INOSINE 5'-TRIPHOSPHATE). Given that both entries share a high degree of structural similarity and also bind nucleotide-like ligands, do they also share the same or similar binding sites?

It should be possible answer that question by looking at the ligand interaction results for these two hetgroups from the corresponding entries. Click on the

"interactions" link for each of the two ligands as shown above. This takes you to the details of the residue interactions in both entries using another PDBe service (PDBeMotif: <u>http://pdbe.org/motif/</u>).

<u>2MJP</u>

hound molecules			
No. Motif/Active site	Ligand	Environment	
PKC PHOSPHO SITE stmotif betatum 1 niche alphabetamotif stturn niche	A 176÷178 A 176÷180 A 149÷152 A 151÷154 A 152÷155 A 151÷155 A 151÷155 A 15÷17 A 15÷17	1만 500A — <mark>ASN 17A GLU 23A SER 89A ASP 152A HIS 177A ARG 178A PHE 115A HOH 588A ASN 19A LYS 20A SER 74A GLY 75A LYS 90A P</mark> H	<u>1E</u> 149A
<u>2E5X</u>			
bound molecules			
No. Motif/Active site Ligan	d Environm	nent	
1 💰 🆧	🏦 <u> </u> 201A 📄 <u>NA 301A </u>	NA 302A THR 7A LYS 12A SER 66A ARG 169A HOH 414A HOH 450A HOH 529A HOH 551A ASN 9A SER 82A PHE 140A GLY 141A TYR 142A SER 8A	ASP 65A

Now going back to the residue-by-residue 3D-mapping results provided by PDBefold, we can make some very interesting observations. For example ASN 17 and LYS 20 interacts with the ligand ANP in 2MJP, aligns with ASN 9 and LYS 12 from PDB entry 2E5X. Both ASN 9 and LYS 12 are also involved in interactions with ITT in a similar manner.

•	A:THR	15	::•	1.83	•	A:THR	7
-	A:GLY	16		2.95	•	A:SER	8
+	A:ASN	17	::•	3.82	H+	A:ASN	9
H+	A:PRO	18	::•	4.87	H+	A:PRO	10
H+	A:ASN	19		5.09	H-	A:GLY	11
H+	A:LYS	20	:::•	3.70	H+	A:LYS	12
H-	A:ILE	21	::	3.77	H-	A:VAL	13
H+	A:LYS	22	::	4.19	H+	A:ARG	14
H+	A:GLU	23	:::	3.13	H+	A:GLU	15
H-	A:ALA	24	:	2.22	H–	A:VAL	16

In this way if we compare the rest of the residues present in the binding environment for ANP and ITT, it will be very clear that both the proteins try to adopt similar binding environment in order to interact with a nucleotide ligand. This observation also indicates that the hypothetical protein represented by the PDB entry 2MJP is potential pyrophosphatase similar to the protein under PDB

EUROPE

entry 2E5X. Therefore in this case, the function of the protein molecule dictates its overall structural fold rather than its sequence identity.

Go back to all the results of the structural comparison of 2MJP with the PDB.

<u>9</u>	0.66	8.2	8.5	2.04	181	3	49	93	2zti:A	100	184	ø	STRUCTURES OF DIMERIC NONSTANDARD NUCLEOTIDE TRIPHOSPHATE PYROPHOSPHATASE FROM PYROCOCCUS HORIKOSHII 073: FUNCTIONAL SIGNIFICANCE OF INTERPROTOMER CONFORMATIONAL CHANGES
<u>10</u>	0.66	6.7	7.7	1.93	177	5	50	79	2dvp:A	92	184	0	STRUCTURE OF NTPASE FROM PYROCCOUS HORIKOSHII
<u>11</u>	0.65	7.9	8.3	2.08	181	3	49	93	2e5x:A	100	185		STRUCTURE OF NUCLEOTIDE TRIPHOSPHATE PYROPHOSPHATASE FROM PYROCOCCUS HORIKOSHII 0T3
12	0.65	8.7	8.8	1.42	168	7	35	71	2car:B	83	194		CRYSTAL STRUCTURE OF HUMAN INOSINE TRIPHOSPHATASE
<u>13</u>	0.64	8.6	8.8	1.48	170	7	35	71	2car:A	83	196		CRYSTAL STRUCTURE OF HUMAN INOSINE TRIPHOSPHATASE
14	0.64	8.8	8.8	1.41	168	7	35	71	2i5d:A	83	195		CRYSTAL STRUCTURE OF HUMAN INOSINE TRIPHOSPHATE PYROPHOSPHATASE
<u>15</u>	0.63	5.4	7.0	1.81	173	6	35	71	lvp2:A	91	189	•	CRYSTAL STRUCTURE OF PUTATIVE XANTHOSINE TRIPHOSPHATE PYROPHOSPHATASE/HAM1 PROTEIN HOMOLOG (TM0159) FROM THERMOTOGA MARITIMA AT 1.78 A RESOLUTION
<u>16</u>	0.63	5.3	7.0	1.87	174	5	34	71	lvp2:B	91	189		CRYSTAL STRUCTURE OF PUTATIVE XANTHOSINE TRIPHOSPHATE PYROPHOSPHATASE/HAM1 PROTEIN HOMOLOG (TM0159) FROM THERMOTOGA MARITIMA AT 1.78 A RESOLUTION
17	0.62	7.9	8.5	1.78	169	8	35	71	2j4e:G	77	186		THE ITP COMPLEX OF HUMAN INOSINE TRIPHOSPHATASE
<u>18</u>	0.61	8.2	8.6	1.76	169	7	34	71	2j4e:E	83	190		THE ITP COMPLEX OF HUMAN INOSINE TRIPHOSPHATASE
<u>19</u>	0.60	3.8	6.8	1.81	177	7	33	71	lk7k:A	91	207	Ξ	CRYSTAL STRUCTURE OF RDGB- INOSINE TRIPHOSPHATE PYROPHOSPHATASE FROM E. COLI
20	0.60	3.4	6.7	1.79	176	6	32	71	2pyu:A	83	208		STRUCTURE OF THE E. COLI INOSINE TRIPHOSPHATE PYROPHOSPHATASE RGDB IN COMPLEX WITH

The checkbox shown on the results page can be used to select entries to submit for multiple alignment from the pairwise alignment pages. Check a few of these results and choose "Submit for Multiple Alignment" from the bottom of the results page. This will do a multiple alignment between the chosen entries and the results will be shown similar to that seen for the pairwise alignment. You may choose the "View Superposed" button to see all your chosen entries aligned.



As can be seen all our chosen entries appear to share the same overall structure.

Multiple Alignments using PDBeFOLD

From the PDBeFOLD submission form, choose the radio button for "Multiple Alignment" to bring up the submission form for multiple alignments.

The multiple alignment option provided by the PDBefold service is very useful when you have more than two proteins for which you need to compare structural folds.

Submission Fo	rm for multiple 3D alignment pairwise explanation of input
List of entries	PDB::1v7r
PDB::1b78:A,B," PDB::2dvn:A,B PDB::2e5xA," PDB::1v7r:A,"	Source: PDB entry PDB code 1v7r view Select chains Find chains Chains: A,"
	Viewer: Jmol 💌
Delete entry	Actualize New entry
Subm	it your query Back to Home Page

In query page for "multiple alignment", type the PDB ID of your choice and click on the button saying "Find chains". This will upload the ID code in the "List of entries" column as shown above. Once an entry is loaded, click on the "new entry" button to upload the next entry in a similar way. The delete entry button enables the user to remove the highlighted PDB ID code from the list of entries.

The **Actualize** button updates data for the highlighted entry from the List of entries, from the details entered in the right part of the Submission Form. Once all the PDB entries are uploaded, click on the "submit your query" button.

		Back to	query	Dov	vnload ×ML	Download to	ext	
##		Structure	N	Neer	Consens	sus scores		
**		Suucture	res	INSSE	RMSD	Q-score		
1		PDB 1b78:A,B,	368	28	1.5246	0.3931	view	download
2	V	PDB 2dvn:A,B	372	26	0.5325	0.4743	view	download
3		PDB 2e5x:A,	185	13	0.6542	0.9391	view	download
4		PDB 1v7r:A,	186	13	0.5231	0.9496	view	download
	0	Number of alig Select all Unsele	ned S ctall se wh	SEs view s	13 Ov superposed ries	download FA	0.31 STA alig	gnment
	det	ails PDB 1b78	econo A,B	dary Si SHSF	tructure A	lignment SHHH shshsh	hsshst	hh
SSE		PDB 2dvn	:A,B	SHSH	ISHHSS-	SHHH	heech	h-
SSE		PDB 2dvn PDB 2e5x	:A,B :A	SHSH	ISHHSS-	SHHH shshsh SHHH	hssshl	h-

The multiple alignment results page shows structural alignment among the 4 entries that were uploaded (1b78, 2dvn, 2e5x and 1v7r). You can also view the superposed entries in a graphics viewer by pressing the "Show Superposed" button.



The central domain region in all proteins appears to be highly similar. Analysis of the residue-by-residue mapping data (similar way it was done in the pairwise alignment before) also indicates high degree of similarity observed in the active binding site (nucleotide binding) of all these proteins.

			3	D Struc	tura	al	ig	nmer	nt				
	1b78			2dvn				2e5x	:A		1v7r	:A	
				A:MET	1			MET	1		MET	1	
	A: LYS	10	8	A: LYS	2	Ш	8	LYS	2	111 🖻	TA 8	2	
s	A:ILE	11	III s	A:ILE	3	Ш	s	ILE	3	III s	ILE	3	
s	A: TYR	12	III s	A: PHE	4	Ш	s	PHE	4	III s	PHE	4	
s	A: PHE	13	S	A: PHE	5	Π	g	PHE	5	III 8	PHE	5	
s	A: ALA	14	S	A:ILE	6	Ш	8	ILE	6	III s	ILE	6	
	A: THR	15	Π	A: THR	7	Π		THR	7		THR	7	
	A: GLY	16	Ш	A: SER	8	Ш		SER	8	Π	SER	8	
н	A: ASN	17	Шн	A: ASN	9	Ш	н	ASN	9	ШН	ASN	9	
н	A: PRO	18	Шн	A: PRO	10	Ш	н	PRO	10	ШН	PRO	10	
н	A: ASN	19	Шн	A: GLY	11	Π	н	GLY	11	ШН	GLY	11	
н	A: LYS	20	Шн	A: LYS	12	Ш	н	LYS	12	ШН	LYS	12	
H	A:ILE	21	ШН	A:VAL	13	Ш	н	VAL	13	ШН	VAL	13	
н	A: LYS	22	III H	A: ARG	14	Ш	н	ARG	14	III H	ARG	14	
н	A: GLU	23	ШН	A: GLU	15	Ш	н	GLU	15	H	GLU	15	
H	A: ALA	24	III H	A:VAL	16	Ш	н	VAL	16	H	VAL	16	
н	A: ASN	25	Шн	A: ALA	17	Ш	н	ALA	17	III H	ALA	17	
н	A:ILE	26	III H	A: ASN	18	Ш	н	ASN	18	ШН	ASN	18	
н	A:ILE	27	ШК	A: PHE	19	Ш	н	PHE	19	H	PHE	19	
н	A:LEU	28	III H	A: LEU	20	Ш	н	LEU	20	H	LEU	20	
	A:LYS	29	н	A: GLY	21	Ш	H	GLY	21	III H	GLY	21	

These results obtained from PDBefold service provide encouraging prospects of understanding possible roles of a hypothetical protein or structural genomics proteins whose function is yet to be determined.

OTHER EXAMPLES

Secondary structure alignments can often show relationships that are not immediately obvious from sequence identity alone. Here are a few example which you may find interesting.

a) Alpha-lactalbumin (PDB entry 1A4V <u>http://pdbe.org/1a4v</u>).

Start a PDBeFOLD comparison for all entries in the PDB archive against 1A4V. Once the results are shown, scroll to the bottom of the page and sort by %seq instead of Q-score.

Sort by	Q-score 🛟	arrange by SCOP family	match 1 jump
0.014	rmsd align length Q-score		
>>>	P-score Z-score Seq %	esults as a plain text file	
>>)	matched SSEs number of gaps	IL file	

Now scroll to the last page and choose one of the results from the last

p	ag	ge.												
6	660	0.76	6.5	7.5	1.45	122	5	34	80	21z2:A	100	129		THE THREE DIMENSIONAL STRUCTURE OF TURKEY EGG WHITE LYSOZYME AT 2.2 ANGSTROMS RESOLUTION
ę	661	0.57	2.9	5.1	2.16	114	6	33	70	2h5z:A	100	122		CRYSTALLOGRAPHIC STRUCTURE OF DIGESTIVE LYSOZYME 1 FROM MUSCA DOMESTICA BOUND TO CHITOTETRAOSE AT 1.92 A RESOLUTION
9	662	0.55	2.9	5.1	2.23	115	5	31	70	3cb7:B	100	126		THE CRYSTALLOGRAPHIC STRUCTURE OF THE DIGESTIVE LYSOZYME 2 FROM MUSCA DOMESTICA AT 1.9 ANG.
	Т	ЭТ		\cap				T	Т		Т	Δ.		DANK ELIDODE
Е	nt	ry	3	cb	7 h	as	31	.%	sec	quer	ice	id	lei	ntity but 70% structure similarity

between the two proteins. Look at the details of the match. The structures are highly similar.

		(Back to r	N natch list	latc						
Qu	iery P	DB 1a4	v:A		Alig	DB					
Nres	%res	N	%SSE	Q	Ρ	RMSD	Nalgn	Nres	%res	N	
123	93	10	70	0.549	2.94	2.233	115	126	91		
				%seq	Z	N _{SSE}	N gaps	THE	E CRYST	ALLO	
ALPHA-LACTALBUMIN 31.3 5.08 7 5 LYSO										2 FRO CA AT	
vie	ew do	ownload		1	view	w) (
				🗉 SU	perp	/iew					
			5	Second	lary						
					la4v: 3cb7:	A HhHs: B H-H	вННННН -ННННН				1-

b) Eosinophil Major Basic Protein (PDB entry 1H8U http://pdbe.org/1h8u).
Do a search for structural similarity for PDB entry 1H8U against the whole PDB archive. Once the results are in, resort the results by %seq identity as in the previous example. Scroll to the last page.

19	1 0.39	5.7	7.4	1.65	85	7	14	88	lkcg:B	70	123		NKG2D IN COMPLEX WITH ULBP3
<u>19</u>	2 0.38	0.6	3.1	2.01	93	8	14	75	lkg0:C	75	136	•	STRUCTURE OF THE EPSTEIN-BARR VIRUS GP42 PROTEIN BOUND TO THE MHC CLASS II RECEPTOR HLA-DR1
19	3 0.43	5.9	7.5	1.89	94	8	14	88	1mpu:A	78	128	8	CRYSTAL STRUCTURE OF THE FREE HUMAN NKG2D IMMUNORECEPTOR

The last hit in the list is PDB entry 1MPU. This entry has 14% sequence identity with our query structure while sharing 88% structural identity. Look at the details of this hit and view the superposed entries as previously.



Scroll down the details page and look at the residue-by-residue listing. All the CYS residues between the two structures are conserved and at the same equivalent positions. These residues form the disulphide bonds in the two structures that keep the scaffold of the protein intact. As a matter of fact, both these proteins belong to a large family of sugar-binding proteins called c-type lectins. All c-type lectins share the same overall structure constrained by the disulphide bonds, and not all proteins in this family actually bind sugars. c) PDB entry 1TIM and 2055. Do a pairwise alignment between 1TIM and 2055 using the pairwise alignment form. The results will show that the two proteins share 4% sequence identity and 70% structural similarity.



Structure Alignment Results

Both proteins have minimal sequence identity and yet belong to the same fold class (the TIM barrel).

This ends our tutorial on PDBeFOLD. We hope you found this useful and will be able to use this tool in your future research and analysis. It is hopefully clear from the examples given in this tutorial that fold space is more restricted than sequence space and most proteins tend to fall broadly into one of the many fold classes already found in the PDB. If you need to get in touch with the PDBe regarding any aspect of the programme, please email <u>pdbehelp@ebi.ac.uk</u> and we will try to assist you in any way possible.